WORLD INTELLECTUAL PROPERTY ORGANIZATION International Bureau



INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(51) International Patent Classification 3:		(11) International Publication Number:	WO 82/ 02387
C08F 2/02	A1	(43) International Publication Date:	22 July 1982 (22.07.82)

(21) International Application Number: PC

PCT/US82/00010

(22) International Filing Date:

7 January 1982 (07.01.82)

(31) Priority Application Number:

223,597

(32) Priority Date:

9 January 1981 (09.01.81)

(33) Priority Country:

US

(71) Applicant: S. C. JOHNSON & SON, INC. [US/US]; 1525 Howe Street, Racine, WI 53403 (US).

(72) Inventors: HAMIELEC, Archie, E.: 99 Rennick Road, Burlington, Ont. L7R-3X5 (CA). LAWLESS, Grant, P.: 4724 Chickory Lane, Racine, WI 53403 (US). SCHULTZ, Harold, H.: 3726 Sherrie Lane, Racine, WI 53405 (US).

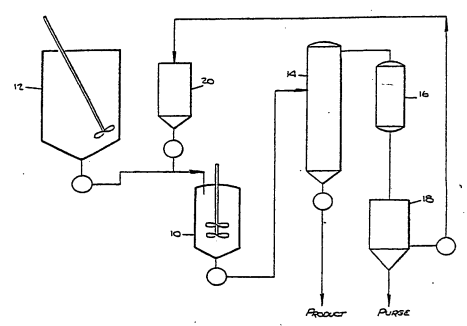
(74) Agents: CHRYSTAL, John, J. et al.; Ladas & Parry, 104 South Michigan Avenue, Chicago, IL 60603 (US).

(81) Designated States: AT (European patent), BE (European patent), CH (European patent), DE (European patent), FR (European patent), GB (European patent), JP, LU (European patent), NL (European patent), SE (European patent).

Published

With international search report.

(54) Title: PROCESS FOR CONTINUOUS BULK COPOLYMERIZATION OF VINYL MONOMERS AND PRODUCT THEREOF



(57) Abstract

A process to bulk polymerize vinylic monomers to prepare low molecular weight polymers employing thermal initiation at reaction temperatures from 235°C to 310°C and residence times of at least 2 minutes in a continuous stirred reactor zone containing the molten resin mix.

FOR THE PURPOSES OF INFORMATION ONLY

Codes used to identify States party to the PCT on the front pages of pamphlets publishing international applications under the PCT.

•		KP	Democratic People's Republic of Korea
ΑT	Austria	LI	Liechtenstein
AU	Australia	ī.K	Sri Lanka
BE.	Beigium	ĬŪ	Luxembourg
32	Brazil	MC	Monaco
CF .	Central African Republic	MG	Madagascar
CG	Congo	MW	Malawi
CH	Switzerland	NL	Netherlands
CM.	Cameroon	NO	Norway
DE	Germany, Federal Republic of	RO	Romania
	Denmark		

. A. 19!

明明	United Kingdom Hungary Japan	TG US	Togo United States of America

PROCESS FOR CONTINUOUS BULK COPOLYMERIZATION OF VINYL MONOMERS AND PRODUCT THEREOF

This invention relates to a process for bulk copolymerizing vinyl monomers by thermal initiation. In particular, it relates to a method for forming low molecular weight polymers of relatively uniform molecular weight by continuous bulk copolymerization.

Styrene and acrylic acid copolymers, conveniently identified as (S/AA), of low molecular weight have found wide acceptance in industry. Such copolymers, sometimes denominated "solution polymers," have been

- 10 employed in inks, floor polishes, paper coatings, paints and adhesives. Typical solution polymers have a number average molecular weight $(M_{\rm n})$ from about 2000 to 5000 and a weight average molecular weight $(M_{\rm w})$ from about 7000 to 10,000. Heretofore, low molecular weight
- 15 solution-type polymers have been produced employing mass polymerization techniques. Typically, by such techniques, vinylic monomers were polymerized in the presence of a solvent and at low temperatures of about 120°C. Often the reaction was expedited with the assistance of cata-
- 20 lytic initiators and chain transfer agents, to help control the molecular weight of the product copolymers. Examples of such polymerization techniques are disclosed in U.S. Patents 2,556,488, 3,673,168, 3,753,958, 3,879,357 and 3,968,059.
- With the cost of petroleum feedstock rising explosively in recent years, the price of vinyl monomers has risen dramatically. Accordingly, there exists a need



Service and the service of the

to reduce the time and expense in preparing low molecular weight yinyl copolymers by simplifying process techniques. The solvents normally employed in solution polymerization are expensive. Solvent recovery equipment must be utilized for economic operation and safety problems have arisen with the use of volatile organic solvents. Similar problems exist when employing chemical initiators or catalysts and/or chain transfer agents in the process. Since solution polymerization is typically carried out in large batches, if the process malfunctions, then an unacceptably large amount of product must be wasted.

Previously, styrene monomer has been homopolymerized to form high molecular weight polymers 15 from 20,000 to 100,000 (M,) in a continuous mass polymerization process without solvents, catalytic initiators and molecular weight regulators, as disclosed in U. S. Patents 2,496,653 and 3,859,268. It has been generally recognized that at temperatures above about 200°C, thermally initiated styrene polymerization produces products having undesired dimers, trimers and oligomers of wide ranging molecular weight and having a high dispersion index (weight average molecular weight -M / number average molecular weight -M l. It has long 25 been desired to produce solution polymers useful in providing an alkali cut on the order of 25 to 35% solids for use in various finishes, polishes and the like or for ink resins. Heretofore, solution polymers at high solids content were unduly viscous and difficult 30 to handle and process. This was due, in part, to their high dispersion index.

In general, for commercial use in solution.

35 of less than about 2. Thermally initiated styrene

BUREAU CMPI

homopolymerization at elevated temperatures has not provided acceptable dispersion indices. In the article by Husain and Hamielec, AICHE Symposium Series, Vol. 72, pp. 112-127 (1976), during bulk thermal polymerization of styrene in a tubular reactor, a "runaway" reaction at 297°C provided a polystyrene with a polydispersibility of 3.68. In Bengouh and Park, European Polymer Journal, Vol. 14, pp. 889-894 (1978), at reactor temperatures of 230°C to 250°C, polydispersity was from 3 to 3.8 after thermal initiated polymerization of styrene monomer.

It has been disclosed in U.S. Patent 4,117,235 that batches of an acrylic monomer can be thermally polymerized in sealed glass tubes at temperatures from 230°-280°C to provide an acrylate polymer with a number average molecular weight of less than about 5000, in the presence or absence of a chain transfer agent or solvent. However, this process is not conducted on a continuous basis. For practical applications, a continuous process is required. Homopolymerization is exemplified. Neither the dispersion index of the resulting material nor its dimer, trimer or oligomer content, was provided.

Accordingly, the art has long sought a

25 continuous bulk polymerization process capable of
selectively producing low molecular weight vinylic
copolymers having a low dispersion index, employing
thermal initiation, without the need of a solvent,
catalyst or weight retarding agent. As employed

30 herein, the term "vinylic polymer" refers to the addition polymer formed by polymerizing vinylic monomers.
Typical vinylic monomers include such diverse monomers
as a-methyl styrene vinyl toluene, styrene, acrylic
or methacrylic acids, acrylic or methacrylic esters
and the like. Further, contemporary industry standards



require that such a process have sufficient flexibility to selectively increase or decrease the molecular weight of the desired product in accordance with market requirements.

According to the present invention, a bulk polymerization process for preparing low molecular weight vinylic polymers having a dispersion index less than about 2 and a number average molecular weight from about 1000 to 6000 comprises the steps of continuously:

- (a) charging a mixture of vinylic monomers into a continuous stirred reactor zone containing a molten resin mixture comprising unreacted vinylic monomers and said vinylic polymers;
- (b) maintaining the molten resin mixture at a reaction temperature from about 235°C to 310°C; and
- (c) maintaining a flow rate through said reaction zone sufficient (1) to provide a residence time of said charged vinylic monomer mixture in said reaction zone of at least about 2 minutes to provide a reaction product; and (2) to maintain a predetermined level of reaction mixture in said reaction zone; wherein said mixture of vinylic monomers comprises at least one monoalkenyl aromatic monomer and at least one acrylic monomer.

In general, a mixture of two or more vinyl type monomers, preferably including styrene and an acrylic monomer, are charged continuously into a reactor zone containing molten resin having the same ratio of vinyl monomers as the monomer mixture. The

Reaction product is pumped out of the reaction zone
35 at the same weight rate as the monomers are charged to



provide a fixed level of resin in the system.

To reduce the unreacted monomer content of the polymer product, the molten resin may be subjected to separation means to remove or reduce the content of unreacted monomer and any volatile by-products. Such unreacted monomer may be recovered and reused in the system.

While applicable to such monoalkenyl aromatic monomers as alpha-methyl styrene, vinyl toluene, para-methyl styrene and tertiary butyl styrene, enhanced results are obtained when the aromatic monomer contains styrene or a mixture of a-methyl styrene and styrene in a weight ratio of from about 2:1:1:2.

A second component of the polymer produced

15 by the present process is an acrylic monomer. The
term "acrylic monomer," as employed herein, includes
acrylic acid alone or mixtures of acrylic acid and an
ester or other drivative of acrylic acid or methacrylic acid. Typical examples of such acrylic acid

20 or methacrylic acid, ester or derivative include:
methacrylate esters, as methyl methacrylate, hydroxy
ethyl methacrylate and dimethylaminoethylmethacrylate;
methacrylic derivatives, such as methacrylic acid and
salts thereof and methacrylonitrile; acrylate esters,

25 such as methyl acrylate and ethyl acrylate; and acrylic
acid derivatives, as acrylonitrile.

Other suitable nonacrylic ethylenic monomers can be added to the monomer charge, including vinyl esters, such as vinyl acetate and maleic acid. The proportions of such monomers can be varied. In certain instances it may be possible to substitute such monomers for the acrylic monomer.

Preferred monomer charges include: the comonomers, styrene and acrylic acid and the termonomers, a-methyl styrene/styrene and acrylic acid. Such



19. Jan 19. 30 m 18 8 18 18

charges provide, respectively, a preferred copolymer (S/AA) and a preferred terpolymer (AMS/S/AA) having a low dispersion index; having low residual monomer content and exhibiting a number average molecular weight adapted to be controlled to a narrow predetermined range by presetting the reaction temperature and residence time in accordance with the invention.

The preferred monomer charge employs from about 60-80% by weight monoalkenyl aromatic monomer and 40-20% by weight of acrylic monomer, such as 10 acrylic acid. Most preferably the monoalkenyl aromatic monomer is styrene or a mixture of styrene and a-methyl styrene in a weight ratio from 2:1 to 1:2. It is particularly preferred to employ 3 parts by weight to 2 parts by weight of styrene solids per part of acrylic acid.

In order to obtain a polymer of the invention with the desired molecular weight and dispersion index to provide a high quality solution-type polymer, the reaction temperature is maintained from about 235°C to about 310°C.

The minimum reaction temperature at steady state will vary, depending on the particular monomers charged to the reactor. Generally, it was found that at temperatures below about 175°C, the material formed was too viscous to process efficiently.

At reaction temperatures below about 235°C the number average molecular weight may exceed the desired maximum of about 6000. Further, the uniformity of the reaction product deteriorates unacceptably. In particular, the dispersion index, an important

In particular, the dispersion index, an important measure of uniformity, shows that the spectrum

when the reaction temperature is maintained below about 235°C.

Enhanced results are obtained and accordingly



it is preferred to employ reaction temperatures from about 245°C to 275°C. Within the preferred temperature range it is possible to achieve the most desirable balance of copolymer properties, such as molecular weight, dispersion index, purity, and ability to form a high solids content, alkali-soluble resin cut.

At temperatures above about 310°C, the temperature can have adverse effects on the product. In addition, at higher reaction temperatures the polymer products tend to be discolored and exhibit undesired yellowing possibly by formation of undesired byproducts, including oxidation products. Further, at such elevated temperatures, the resulting polymer products may be subject to depolymerization, reversible reactions and other side reactions which yield dimers, trimers and other undesired low molecular weight oligomers. Such by-products contaminate resin cuts of product polymer and contribute to their high optical density. This, in turn, indicates that off-color finishes can be expected employing such resin cuts.

In general, the reaction time or residence time in the reaction zone is controlled by the rate of flow of constituents through the reaction system. The residence time is inversely proportional to flow rate. It has been found that at a given temperature, the molecular weight of the polymer product decreases as the residence time increases.

In accordance with these factors it is therefore preferred to utilize reaction residence times of at least about 2 minutes to provide satisfactory reaction completion. While the residence time in the reaction zone may be as long as 1 1/2 hours, normally discoloring reactions and other side reactions will dictate that shorter residence times be employed. For most cases a residence time of from about 5 to 45



minutes, and, preferably, from 10 to 20 minutes is satisfactory. In general, even longer residence times increase the yield of product, but the rate of increase of product is generally very slow after about 20 to 30 minutes of reaction.

The flow rate of reactants for the preferred styrene/acrylic acid monomer mix is between about 0.1 to 1.5 pound per minute per gallon of reactor capacity. The flow rate selected for a particular monomer system will depend upon the reaction temperature, constituents, desired molecular weight, desired dispersion index and the particular system employed.

For best results, to produce a given resin of a desired $M_{\rm n}$ and $M_{\rm w}$ with low residual monomer, the reaction temperature and residence times are mutually manipulated in accordance with the principles provided herein.

is a function of residual vapor pressure of unreacted monomer and other volatiles present either in the feed, such as water, or in the reaction mix (produced in side reactions). Although at steady state the process is carried out under a positive pressure, the reaction pressure has no significant effect on the yield and the upper limit is a function of equipment capability, while the lower limit is a function of feed rate and monomer composition. At higher temperatures, the higher gas pressures which result may require special equipment and procedures for safe handling.

In a preferred embodiment of the invention a solvent, capable of forming a reaction product with

cosity of the product. The solvent also acts to lower the viscosity of an alkali cut of the copolymer at high solids contents from about 25 to 35% non-volatiles.



It has been found that a portion of the solvent is strongly associated (by reaction or binding) with the copolymer, while the remaining portion is loosely associated with the copolymer.

For best results from about 1 to 20% by weight of vinylic monomers of the solvent, is employed. While greater or lesser amounts can sometimes be employed, there is an insufficient reduction in viscosity, when less than about 1% of solvent is employed. When 10 amounts greater than about 20% by weight of solvent are utilized, then the process tends to become solvent dependent as a typical solution polymerization process. For best results from about 1 to 10% by weight of solvent is preferably employed.

The solvents employed include polyalkoxy mon-15 alkanols. Such polyethers are well known to the art. Typical examples include the family of solvents derived from 2-ethoxyethanol, known to the art as Cellosolve. Illustrative solvents are butyl Cellosolve and Cel-20 losolve acetate. A particularly preferred solvent is diethylene glycol monoethyl ether, known industrially as Carbitol. Other useful solvents include ethylene glycol monomethyl ether and di-ethylene glycol monomethyl ether.

When other solvents, such as methyl isobutyl ketone (MIBK), are employed the desired results have not been achieved. MIBK does not become associated with the copolymer. When ethylene, propylene or diethylene glycol are employed as the solvent or 30 part of the solvent, the molecular weight distribution will be widened or gelation can occur depending. on the level of the dihydric alcohol and the degree of cross-linking which occurs.

It has been found that the addition of 35 solvent reduces the acid number of the polymer product.



More importantly, it also permits formulation of an alkali soluble resin cut having a solids content from about 25 to 35% non-volatiles at acceptable viscosity.

A preferred embodiment of the process of 5 the present invention is illustrated in the accompanying drawing, which is a flow sheet showing the polymerization process.

The process of the present invention involves the use of a variable fillage type stirred re-10 actor for the polymerization of vinylic monomers to copolymers having a narrow-molecular weight distribution by the proper balancing of the conditions of polymerization and flow rates. The reaction zone as represented by reactor 10 can comprise a continuous 15 stirred tank reactor of any type adapted for variable fillage operation of from as low as 10% to 100% of the usable volume thereof for the production of vinylic polymers. This continuous stirred tank reactor may be either horizontal or vertical and should have provision for close control of the temperature therein by any desired means, including control by cooling jacket, internal cooling coils or by withdrawal of vaporized monomer followed by condensation thereof and return of the condensed monomer to the reaction zone. It will be apparent to those skilled in the art that this reaction zone can, if desired, be constituted by a plurality of continuous stirred tank reactors operated in series. Likewise, it will be apparent that such reaction zone can comprise more than one continuous stirred tank reactor operated in parallel if it is desired to employ several relative-

reaction zone rather than a single large reactor for that purpose.

A preferred form of continuous stirred re-



actor which has been found quite suitable for carrying out the process is that general type illustrated in the accompanying drawing wherein a tank reactor is provided with cooling coils sufficient to remove any 5 heat of polymerization not taken up by raising the temperature of the continuously charged monomer composition so as to maintain a preselected desired temperature for polymerization therein. Preferably also such continuously stirred tank reactor will be pro-10 vided with at least one and usually more vaned agitators driven by an external power source such as a motor. At least one such agitator is positioned to provide agitation of liquid contained in the reactor while operating at minimum fillage, i.e., as low as 15 10% of the volume thereof. Such a continuous stirred tank reactor can, if desired, be provided with additional features for improved efficiency of operation and safety, such as an additional series of internal cooling coils adapted to effectively prevent 20 any "run-away" polymerization if the normal holding period has to be extended for some reason and an outer jacket for additional cooling or heating of the contents of the reactor.

In operation of the present continuous bulk,

25 polymerization process, flexibility and range of choice
can be realized in polymer types produced, as well
as the production rate thereof, by proper choice of
polymerization reaction conditions. The drawing illustrates the operation of the reaction zone of the

30 present process and the manner in which such operation is incorporated into an overall polymer production line. In operation a monomer composition comprising polyalkenyl aromatic and acrylate monomers as
described above, is charged to reactor 10 and the temperature of the monomer feed is raised to from about
235°C to 310°C to produce thermal polymerization.



Reactor 10 is charged from stirred feed tank 12 which contains the reaction charge of monomers. Pressure in reactor 10 can vary from 40 to 400 psia or even higher. It is preferred to operate reactor 10 at from about 50 to about 300 psia, and most preferably from about 150 to about 200 psia.

After the initial fillage of reactor 10 to the desired preselected level and polymerizing the charged monomer to approximately the desired solids content, then the volume of monomer composition charged thereto is adjusted to a value to maintain such preselected level of liquid in reactor 10. Thereafter, the liquid mixture of polymer and monomer is withdrawn from reactor 10, to maintain the preselected 15 level of such liquid mixture in the reaction zone. Polymerization conditions are continuously maintained in reactor 10 to produce a polymer of selected molecular weight and selected degree of conversion or weight percent solids of polymer in such liquid mix-The reaction zone can be operated so as to produce a liquid mixture with a polymer concentration or percent solids from as low as 30 percent to as high as 95 percent by weight and such polymer can possess a number average molecular weight ranging from 1000 to 6000. The level of fillage of reactor 10 can vary from as low as 10 percent to as high as 100 percent, usable volume, and may be controlled by any desired means, for example, a level controller and associated valve or pump in the transfer line from reactor 10. 30

Any desired means of controlling the temperature within reactor 10 may be employed. It is

ternal cooling coils in those reactors so equipped such as reactors of the type illustrated. The entry



of relatively cool monomer composition serves to remove the greater proportion of the heat of polymerization released and the internal cooling coils serve to remove the remainder so as to control the temperature of the liquid mixture therein to a preselected value and thus produce a polymer of the desired degree of conversion and average molecular weight.

As the polymer concentration increases, the possibility of damage from "run-away" reactions is substantially reduced. In general it is preferred to produce in the reaction zone a solids content of from 80 to 95 percent by weight of a polymer of relatively average molecular weight of from about 1000 to 6000 and of relatively narrow molecular weight distribution. The residence time in the reaction zone can vary from about 5 to 45 minutes.

In the drawing there is illustrated a zone of devolatilization. In one embodiment devolatizer 14 is a stirred tank adapted to received unreacted mono-20 mers and polymer product at atmospheric conditions. The process of the present invention can be operated by the use of a single or multiple zones of devolatilization, as desired. In the process outlined the vaporized monomers as well as any low oligomers thereof, are 25 removed from the devolatilization zone, liquefied in a condensor 16 and passed to a receiver 18. From the receiver a stream of the condensed monomers and oligomers can be recycled to recycle tank 20 and then . to reactor 10 as shown. If desired, the oligomers can 30 be vaporized in the devolatilization zone and separated from the vaporized monomers prior to their condensation. Such oligomers can be separately recycled to a reaction zone or purged from the process. In general, the equipment for the present process is known to the 35 art and has been described for use in other bulk polymerization procedures in U.S. Patent No. 3,968,059 and



- 14 -

U.S. Patent 3,859,268.

The resin product pumped from the devolatizer 14 can be solidified by appropriate means or cut into an appropriate solvent system. The resin product may 5 be solidified by employing a conventional flaker. The product flakes can be packaged in accordance with known techniques. For example, the flakes may be conducted to a flake bin by suction and then transported to a bagger.

10 The following examples are set forth to illustrate more clearly the principles and practice of this invention to one skilled in the art. As such, they are not intended to limit the invention but are illustrative of certain preferred embodiments. In these Examples molecular weight of polymer product was determined in accordance with conventional gel permeation chromatography.



30

- 1:5 -

Example I

A vertical stirred tank reactor of 1 gallon capacity adapted for cooling by oil was filled to 50% of its volume from a feed tank containing a mixture 5 of 23 parts alpha-methylstyrene, 45 parts styrene and 32 parts acrylic acid and brought to a polymerization temperature of 272°C. Polymerization was begun with stirring and continued until the solids content of the liquid reaction mix reached about 90% solids. Thereafter, a feed rate of 0.80 pounds per minute of the aforesaid monomer mixture per gallon of reactor was maintained and a residence time of 5 minutes was attained in the first reactor. The temperature of the polymerizing liquid in the first reactor was main-15 tained at 272°C by circulating oil through the reactor jacket. A continuous withdrawal of polymer and monomer was established to maintain the initial level in the reactor.

A stirred devolatizer tank was filled with

the effluent from the reactor to 10-20% of its volume.

The temperature in the devolatized tank was maintained at 272°C. Vaporized monomers and oligomers were condensed and monomer mix was recylced to a recycle feed for delivery to the tank reactor.

A terpolymer product was recovered from the tank having a number average molecular weight of 2640, a weight average molecular weight of 4500, a dispersion index of 1.70 and an acid number of 190. The terpolymer product was processed into flakes.

The flakes were processed into a 20% non-volatiles, resin cut according to conventional procedures. The optical density of the resin cut was 0.1 at 400 nanometers (nm).



- 16 -

Example II

In order to demonstrate the effect of reaction temperature on the molecular weight, dispersion index and optical density of a 20% product resin cut, test runs were conducted in accordance with the procedure set forth in Example I. For each run, the monomer charge, reaction temperature, T, (in °C), feed rate (lbs./min.), residence time (R/T) (in minutes), number average molecular weight Mn of product, weight average molecular weight Mw of product, dispersion index Mw/Mn of product, and optical density, O.D., (log IO) of product were measured. The results are set forth in Table 1.



Table 1

	(mu 00							-1	7 –					•			
	O.D. (400 nm)	5+	7	0.95	0.42	0.41	0.18	0.19	0.16	0.11	0.18	0.07	0.11	0.08	0.04	90.0	0.25
	Mw/Mn	1.36	1.39	1.97	19.1	. 1.74	1,57	1.98	1.94	1.66	1.90	1.98	1.74	1.85	1.68	1,91	2.55
	Mw	1770	1540	2400	2900	3600	3140	5300	5500	5200	6500	7100	6 200	8600	4900	10,600	38,100
	Mn	1300	1110	1630	1800	2070	2000	2680	2830	3130	3430	3580	3570	4660	2920	5560	14,920
Temp.	ပ္စ	309	300	297	289	287	278	276	271	267	263	260	254	249	246	238	223
Residence time	minutes	10	20	10	. 10	10	15	10	10	10	10	. 01	រ ភ	ហ	1.5	រភ	10
Feed Rate	1bs/min	0.4	0.2	0.4	ı	0.4	0.26	0.4	0.4	0.4	0.4	0.26	0.80	0.80	0.26	08.0	0.4
Monomers	parts/parts	68S*/32AA**	37AMS/31S/32AA	68S/32AA	72S/28AA	68S/32AA	68S/32AA + 10% DEGMEE	68S/32AA	68S/32AA	68S/32AA	68S/32AA	72S/28AA	39AMS***/33S/28AA 0.80	24AMS/48S/28AA . 10% DEGMEE	37AMS/31S/32AA	39AMS/33S/28AA	68S/32F.A

*** AMS = a-methylstyrene ** AA = acrylic acid * S = styrene

• DEGMEE = diethylene glycol monoethyl ether

The results illustrate that the reaction temperature is an important factor in the present process. When reaction temperatures are set below about 235°C, the molecular weight of the polymer product is greater than 6000 and the dispersion index becomes greater than 2. Similarly, as reaction temperatures rise above 310°C, the molecular weight of the product is reduced to the high oligomer range and the optical density is severely compromised. The lower the value for optical density, the clearer the polymer product.

Example III

In order to evaluate the effect of residence time on the process a series of test runs were con15 ducted and reported in accordance with the procedure of Example I and the results were recorded in Table 2.



		Tal	Table 2				
Monomers parts/parts	Feed Rate 1bs/min	Reaction temp.	Residence time minutes	Mn	MM	0.D. (400 nm)	
68S/32AA + 10% DEGMEE	1.16	253	1.0	5640	е	0.03	
31S/37AMS/32AA	.1.0	274	4	2370	1.67	0.81	
68S/32AA + 10% DEGMEE	0.8	278	ហ	2780	1.72	60.0	
68S/32AA + 10% DEGMEE	0.4	263	G	3060	1.76	0.07	
68S/32AA	0.4	267	10	3130	1.66	0.11	
68S/32AA + 10% DEGMEE	0.26	278	1.5	2000	1.57	0.18	
37AMS/31S/32AA	0.23	260	20	2010	1.59	. 0.28	
68S/32AA	0,23	266	23 .	2590	1.62	0.18	
68S/32AA + 10% DEGMEE	0.23	261	25	2690	1.67	0.11	
68S/32AA + 10% DEGMEE	0.18	260	45	2520	1.67	0.13	
68S/32AA	0.12	266	50	2150	1.57	0.37	



- 20 -

The results show the significant effect of residence time of reaction mixture on the process.

As the residence time is reduced to less than about two minutes, the molecular weight and dispersion index are adversely effected.

Example IV

In order to illustrate the effects of employing a solvent in the monomers mix, tests were run in accordance with the procedure of Example I. The quantity of solvent added to the monomers mix was varied between 5 and 20% by weight of total monomers mix. For illustrative purposes a test run is provided wherein no solvent was employed. The results are reported in Table 3. Mn is number average molecular weight and Mw is the weight average molecular weight. The acid number is the number of milligrams of potassium hydroxide required to neutralize one gram of the substance.



Table 3

Monomer	*DEGMEE*	T(°C)	Res. Time (min.)	Feed Rate (1bs/min)	Mu	Mw	Optical Density (400 nm)	Acid
72S/28AA	ហ	277	10	0.5	2770	1.81	0.15	
72S/28AA	• •	274	. 01	0.4	2450	1.64	0.15	
72S/28AA	10	263	15	0.26	3370	1.75	0.10	
72S/28AA	Ö	. 797	æ	0.80	3140	1.72	0.10	
37ams/31s/32aa	50	263	28	0.18	1700	1.56	0.26	
37AMS/31S/32AA	0	. 097	20	0.23	1820	1.54	0.42	
								-

*DEGMEE is diethylene glycol monoethyl ether.

BUREAU

- 22 -

As is evident from the results tabulated hereinabove, the use of a solvent can vary the acid number of the product.

Example V

5 The temperature dependency of the present process when a solvent is additionally employed is illustrated in the tabulation of test runs conducted according to Example 1, set forth in Table 4.



	• .0.0	2.0	0.05	.07	.03	.21	.19	0.16	.05	. 28
•	Mw	1.43	1.64	1.90	1.78	1.67	1.94	1.92	2.33	2.46
	Mn	1120	2200	3570	3380	3650	4530	4690	6580	10,270
	Reaction Temp.	299	262	257	252	249	243	241	221	178
Table 4	Feed Rate 1bs/min	0.2	0.26	0.8	0.4	0.4	0.4	0.4	0.4	1.3
	Reaction Time Minutes	20	15	ທຸ	10	10	10	10	10	-
,	* DEGMEE	10	10	10	20	. 20	10	10	70	10
	Monomer	37AMS/31S/32AA	37AMS/31S/32AA	37AMS/31S/32AA	37AMS/31S/32AA	37AMS/31S/32AA	35AMS/30S/34AA	37AMS/31S/32AA	36AMS/30S/34AA	37AMS/31S/32AA

* a 400 nm



والمنبي وكساعو عكما كالأعلة لتعريف العادات والرازان

lo

- 24 -

The polymers produced by the process of the invention can be employed in conventional floor finishing compositions, inks and the like. Other uses will be apparent to those with ordinary skill in this field.

Although the invention has been described in detail by referring to certain preferred embodiments, it will be understood that various modifications can be made within the spirit and scope of the invention. The invention is not to be limited except as set forth in the following claims.



- 25 -

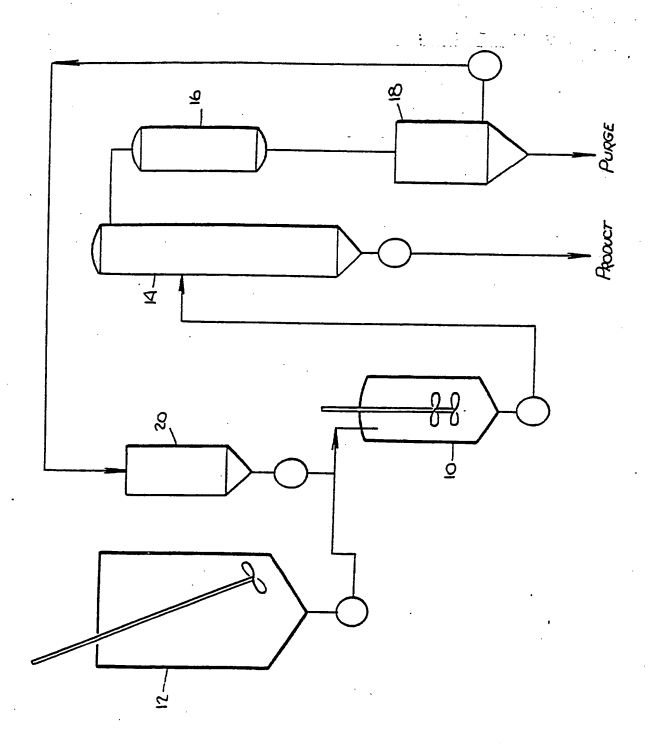
CLAIMS

- 1. Bulk polymerization process for preparing low molecular weight vinylic polymers having a dispersion index less than about 2 and a number average molecular weight from about 1000 to 6000 comprising the steps of continuously:
- (a) charging a mixture of vinylic monomers into a continuous stirred reactor zone containing a molten resin mixture of unreacted vinylic monomers and said vinylic polymers;
- (b) maintaining the molten resin mixture at a reaction temperature from about 235° C to 310° C; and
- (c) maintaining a flow rate through said reaction zone sufficient (1) to provide a residence time of said charged vinylic monomer mixture in said reaction zone of at least about 2 minutes to provide a reaction product; and (2) to maintain a predetermined level of reaction mixture in said reaction zone; wherein said mixture of vinylic monomers comprises at least one monoalkenyl aromatic monomer and at least one acrylic monomer.
- 2. A process according to Claim 1, wherein the monoalkenyl aromatic monomer is from about 60-80% by weight and the acrylic monomer is from about 40-20% by weight.
- 3. A process according to Claim 1, including the step of separating the copolymer reaction product from unreacted monomers and by-products.
- 4. A process according to Claim 2, wherein the monoalkenyl aromatic monomer is styrene or a mixture of a methyl styrene and styrene having a weight ratio from about 1:2 to 2:1 and the acrylic monomer is acrylic acid.
 - 5. A process according to Claim 1, 2,3 or



- 4 wherein the mixture of vinyl monomers includes from about 1-10% by weight diethylene glycol monoethyl ether.
- 6. A process according to Claim 1, 2, 3, or 4, wherein the reaction temperature is from about 245°C to 275°C and the residence time is from about 5 to 45 minutes.
- 7. Bulk vinylic polymers having a dispersion index less than about 2 and a number average molecular weight from about 1000 to 6000 comprising at least one monoalkenyl aromatic monomer and at least one acrylic monomer, the bulk vinyl polymers being prepared by continuously charging a mixture of vinylic monomers into a continuous stirred reactor zone containing a molten resin mixture of unreacted vinylic monomers and said vinylic polymers, maintaining the molten resin mixture at a reaction temperature from about 235°C up to 310°C, and maintaining a flow rate through said reaction zone sufficient to provide a residence time of said charged vinylic monomer mixture in said reaction zone of at least about 2 minutes to provide a reaction product, and to maintain a predetermined level of reaction mixture in said reaction zone.
- 8. Bulk polymers according to Claim 7, wherein the monoalkenyl aromatic monomer is styrene or a mixture of a methyl styrene and styrene having a weight ratio from about 1:2 to 2:1 and the acrylic monomer is acrylic acid.
- 9. Bulk polymers according to Claim 7 or 8 wherein the mixture of vinyl monomers includes from about 1-10% by weight diethylene glycol monoethyl ether.







INTERNATIONAL SEARCH REPORT

International Application No PCT/US 6 2 / 0 0 0 1 C

	OF SUBJECT MATTER (if several classification symbols apply, indicate all) *									
According to Internation	According to International Patent Classification (IPC) or to both National Classification and IPC									
IPC 3	Co8F 2/02									
IL FIELDS SEARCH	10									
Minimum Documentation Searched 4										
Classification System	Classification Symbols									
บร	526/88 209 317 346									
	Documentation Searched other than Minimum Documentation to the Extent that such Documents are included in the Fields Searched 6									

Category •	Citation of C	DERED TO BE RELEVANT 14 Document, 14 with indication, where appropriate, of the relevant passages 17	Relevant to Claim No. 18
x		859,517 Published 25 January 1961	1-6
х	US, A,	4,117,235 Published 26 September 1978	1-6
x	US, A,	4,195,169 Published 25 March 1980	1-6
х	US, A.,	3,035,033 Published 15 May 1962	1-6
A	US, A,	2,556,488 Published 12 June 1951	1-9
Α	US, A,	3,673,168 Published 27 June 1972	1-9
.]			
.			

- Special categories of cited documents: 15
- "A" document defining the general state of the art
- earlier document but published on or after the international filing date
- "L" document cited for special reason other than those referred to in the other categories
- "P" document published prior to the international filing data but on or after the priority date claimed
 "T" later document published on or after the international filing date or priority date and not in conflict with the application, but cited to understand the principle or theory underlying the invention

Date of the Actual Completion of the International Search 2	Date of Mailing of this International Search Report 2
International Searching Authority 5	Signature of Authorized Officer 10 Smith
ISA/US	Edward J. Smith

Form PCT/ISA/210 (second sheet) (October 1977)

International Application No.PCT/US 8 2 / 0 0 0 1 0

FURTHE	R INFORMATION CONTINUED FROM THE SECOND SHEET	
Х	US, A, 3,753,958 Published 21 August 1973	7-9
Х	US, A, 3,879,357 Published 22 April 1975	7-9
A	Aiche Symposium Series, Vol. 72, No. 160, issued 1976, pages 112-127	1-9
A	Bengouh and Park, European Polymer Journal, Vol. 14, issued 1978, pages 889-894	1-9
	- •	
		· ·
	SERVATIONS WHERE CERTAIN CLAIMS WERE FOUND UNSEARCHABLE 19	
	national search report has not been established in respect of certain claims under Article 17(2) (a) for	<u>-</u>
11 Cm	n numbers because they relate to subject matter 12 not required to be searched by this Auti	ority, namely:
l		
ł		
	n numbers, because they relate to parts of the international application that do not comply wi	th the prescribed require.
	to such an extent that no meaningful international search can be carried out 13, specifically:	ni nia biancipad tadma-
}	•	
		•
}		
]		
)		
Ar O	Servations where unity of invention is lacking 11	
This inter	astional Searching Authority found multiple inventions in this international application as follows:	
i ·		
1₽ Asi	Il required additional search fees were timely paid by the applicant, this international search report coveriences application.	rers all searchable claims
2	mly some of the required additional search fees were timely paid by the applicant, this international s	search report covers only
thos	a claims of the international application for which fees were paid, specifically claims:	
Ī	·	ı
	equired additional search fees were timely paid by the applicant. Consequently, this international sear evention first mentioned in the claims; it is covered by claim numbers:	ch report is restricted to
.·		
		•
Remark or		
=	edditional search fees were accompanied by applicant's protest.	•
[∐ ™ [rotest accompanied the payment of additional search fees.	

Form PCT/ISA/210 (supplemental sheet (2)) (October 1977)

•